

U.S. EPA's use of Read-Across in **Provisional Peer-Reviewed Toxicity Value Assessments**

Jason C. Lambert, PhD, DABT U.S. EPA, National Center for Computational Toxicology

Understanding and Applying Read-Across for Human Health Risk Assessment Workshop

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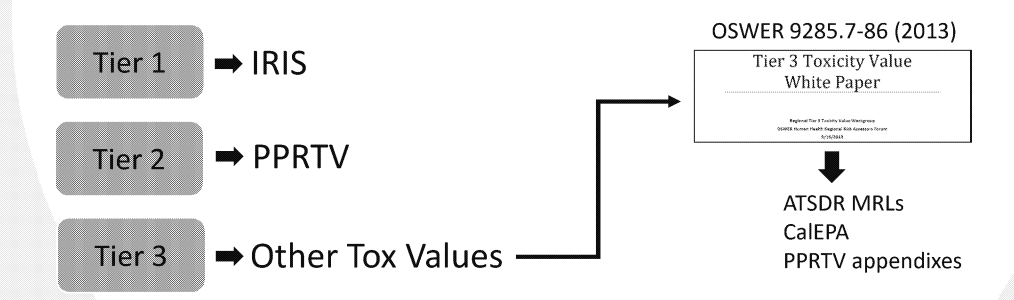
Outline of this presentation

- What are PPRTVs?
- Advent of Appendix Screening Values: Expert-Driven Read Across
- *n*-Heptane example
- p,p'-DDD example (integrating ToxCast bioactivity)
- Limitations and challenges



Provisional Peer-Reviewed Toxicity Values

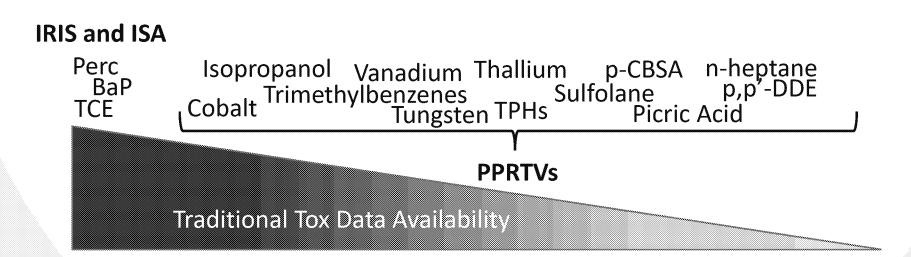
- Annual goal: Derive provisional human health reference values for 12 priority chemicals for OLEM/Regions when such values are not available from IRIS
- OSWER Directive 9285.7-53 (Dec. 5, 2003) established a hierarchy for selecting Human Health Toxicity Values for use in Superfund Risk Assessments





Provisional Peer-Reviewed Toxicity Values

- For over two decades, the PPRTV program has developed human health assessments for chemicals with highly variable hazard databases
- Includes development of subchronic and chronic non-cancer reference values (RfVs)
 and cancer values for chemicals of interest to OLEM/Regions
- A data-rich PPRTV assessment might provide up to six provisional values
- PPRTV 'appendixes' were a key development circa 2007





Expert-driven Read Across

- Approach is based on evidence integration and synthesis to select the best analogue chemical based on three proposed similarity contexts/categories (Wang et al., 2012)
- Similarity Contexts are:
 - (1) structural (including physchem),
 - (2) metabolic, and
 - (3) toxicity-like

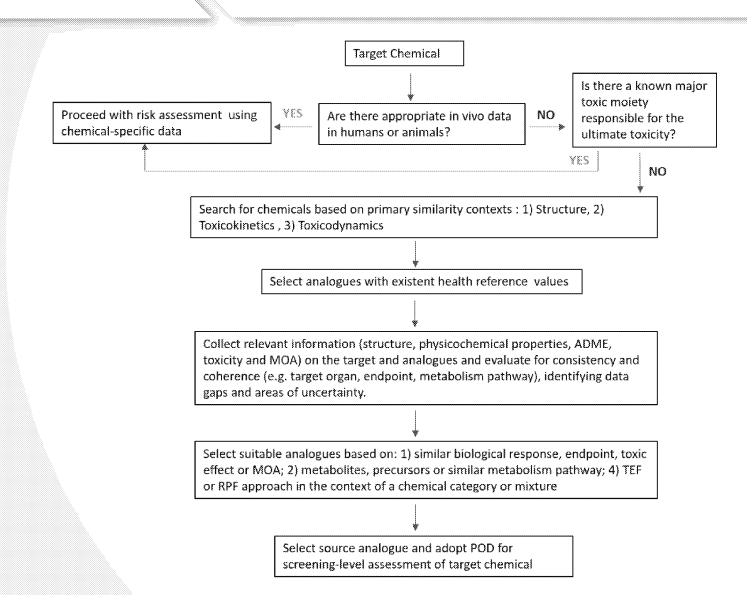


http://www.sciencedirect.com/science/article/pii/S0273230012000323

- Analogue chemicals are selected based on evidence across all three similarity contexts (i.e., analyses are integrated not sequential/linear)
- The POD(s) from the selected analogue is used as a surrogate to derive screening PPRTVs for a data-poor target chemical of concern



General Expert-Driven Read Across Workflow



Adapted from Wang et al., (2012)



n-Heptane (CASRN 142-82-5)



- Saturated aliphatichydrocarbon
- Used as a non-polar solvent and also as a standard in gasoline engine testing
- Requested for human health assessment
- Poor hazard and doseresponse database

Physicochemical Properties of n-Heptane			
Property (unit)	Value		
Physical state	Liquid		
Molecular weight (g/mol)	100.21		
Vapor pressure (mmHg at 25°C)	46		
Henry's Law constant (atm-m³/mol at 25°C) (estimated)	2.27		
Solubility in water (g/L at 25 C)	0.0034		
Octanol-water partition constant (log K _{ow})	4.66		



Identify Structural Analogues of n-Heptane

• Identify commonalities in structural/physicochemical properties between potential surrogates that have available toxicity values, and target chemical of concern

Chemical	n-Heptane	n-Hexane	n-Nonane
Structure			
CASRN	142-82-5	110-54-3	111-84-2
Molecular weight	100.21	86.18	128.26
DSSTox similarity score (%) ^b	100	85.7	87 .5
ChemIDplus similarity score (%) ^c	100	82.7	84.6
Melting point (°C)	-90.6	-95.3	-53.5
Boiling point (°C)	98.5	68.7	150.8
Vapor pressure (mm Hg at 25°C)	46	151.3	4.45
Henry's law constant (atm-m³/mole at 25°C)	2.27 (estimated) ^a	1.71 (estimated) ^a	4 (estimated) ^a
Water solubility (mg/L)	3.4	9.5	0.22
Log K _{ow}	4.66	3.9	5.65
pKa	NA	NA	NA

^{*}Data was gathered from PHYSPROP database for each respective compound unless otherwise specified (<u>U.S. EPA, 2012b</u>). bDSSTox (2015).

NA = not applicable.

^{&#}x27;ChemIDplus Advanced, similarity scores (ChemIDplus, 2016).



Identify Toxicokinetic Similarities

• Identify commonalities primarily in metabolism (metabolic precursors, metabolites), but also absorption, distribution, and excretion, between potential analogue chemicals that have available toxicity values, and target chemical of concern

Table A-2. Summary of Metabolites for <i>n-</i> Heptane (CASRN 142-82-5) and Structural Analogs				
Chemical	Route	Species	Metabolites in Urine	References
n-Heptane	Inhalation (1,800 ppm for 6 h)	Rat/M	2-heptanol (46.3), 3-heptanol (35.2), y-valerolactone (11.5), 2-heptanone (3.5), 3-heptanone (1.5), and 4-heptanone (1.2), 2,5-heptanedione (0.8) over 24 h ^a	Perbellini et al. (1986)
n-Hexane	Inhalation (1,000 ppm for 8 h)	Rat/M	2-hexanol (57), 2,5-hexanedione (33), 3-hexanol (6), and 1-hexanol (3), 2-hexanone (1) over 24 h ^a	Fedtke and Bolt (1986)
n-Nonane	Oral (800 mg/kg-d)	Rat/M	y-valerolactone (38.6), 2-nonanol (17.9), 3-nonanol (10.7), 4-nonanone (6.8), δ -heptanolactone (6.5), 1-heptanol (5.7), 4-nonanol (3.5), 5-methyl-2-(3-oxobutyl) furan (3.2), δ -hexanolactone (2.8), 2,5-hexanedione (1) over 48 h ^b	<u>Serve et al.</u> (1995)

^aPercentage of total metabolites in urine.

^bRelative abundance of metabolites in urine.



Identify Toxicity/Bioactivity Similarities

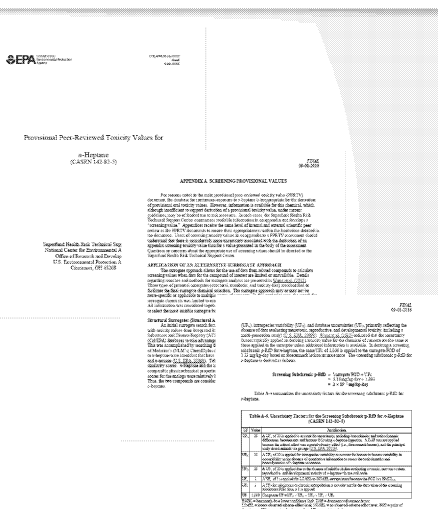
 Identify commonalities in toxicity (e.g., effect levels, target organs/tissues) and/or bioactivity between potential analogue(s) (with available toxicity values) and data-poor target chemical of concern.

Chemical	n-Heptane	n-Hexane	n-Nonane
Structure			
CASRN	142-82-5	110-54-3	111-84-2
Repeat-dose toxici	ty—oral, subchronic		
POD (mg/kg-d)	NA	785	3.13
POD type	NA	LOAEL	BMDL ₁₀
Subchronic UF _C	NA	3,000	1,000
Subchronic p-RfD (mg/kg-d)	NA	3 × 10 ⁻¹	3 × 10 ⁻³
Critical effects	Irritative and proliferative forestomach lesions and potential effects in the liver, kidney, and adrenal glands at a dose of 2,860 mg/kg-d. Lack of neurotoxicity at doses up to 2,860 mg/kg-d based on histological evaluation (13-wk rat study).	Decreased MNCV associated with peripheral neuropathy	Proliferative forestomach esions with varying degrees of hyperplasia and hyperkeratosis of the squamous epithelium
Other effects		Hind-limb paralysis accompanied by evidence of peripheral neuropathy and testicular effects based on histopathology at a dose of 2,843 mg/kg-d (90-d rat study)	Additional effects in principal study: histopathological lesions in the duodenum (rats) and rectum (rats and mice) at doses ≥1,000 mg/kg-d; nasal and pulmonary lesions, possibly due to aspiration (rats and mice). Increases in liver and lung weights at a dose of 5,000 mg/kg-d and dose-related increases in adrenal gland and ovary weights at doses ≥1,000 mg/kg-d. No significant neurohistopathology or neurobehavioral abnormalities reported in rats or mice at doses uto 5,000 mg/kg-d
Species	NA	Rat (M)	Mouse (M) and Rat (F)
Duration	NA	8 wk	90 d
Route (method)	NA	Oral (gavage)	Oral (gavage)



Evidence Integration and Synthesis: n-Nonane as selected analogue

- Similarity Context 1: high structural similarity to n-Heptane (>84%)
- Similarity Context 2: n-Nonane is metabolized $in \ vivo$ similarly to n-Heptane (higher relative amounts of the 2- and 3-alcohol and γ -valerolactone metabolites formed, compared to the neurotoxic γ -diketone compounds from n-Hexane candidate analogue)
- Similarity Context 3: *n*-Nonane-induced proliferative forestomach lesions are similar to the lesions observed after oral *n*-Heptane exposure (as compared to unique *n*-Hexane-induced neurotoxicity)



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Identify Structural Analogues of p,p'-DDD

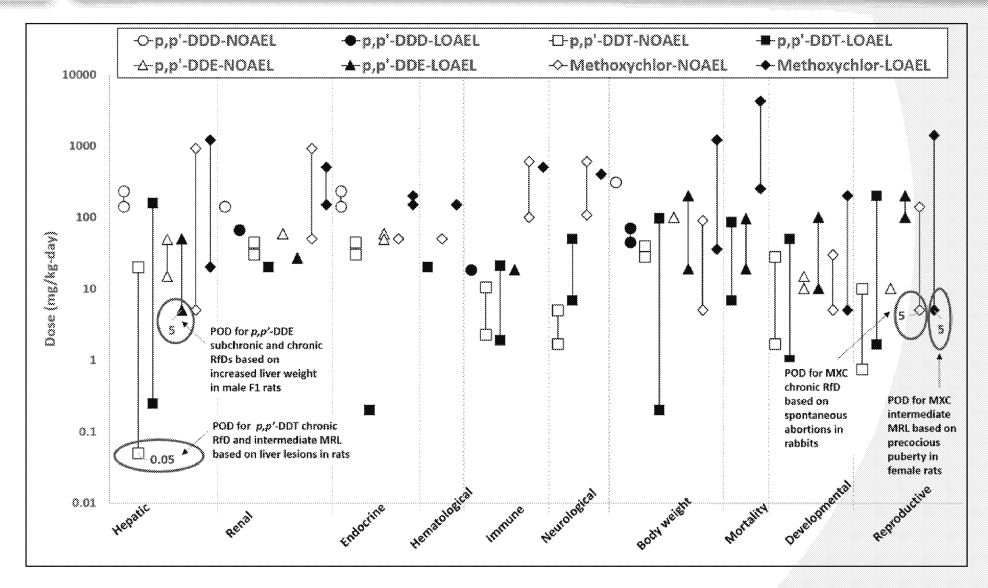
Table 1. Structural Analogues of p,p'-DDD				
	Target Chemical		Analogues ^a	
Name	p,p'-Dichlorodiphenyl dichloroethane (p,p'-DDD)	p,p'-Dichlorodiphenyl trichloroethane (p,p'-DDT)	p,p'-Dichlorodiphenyl dichloroethylene (p,p'-DDE)	p,p'-Dimethoxydiphenyl trichloroethane (Methoxychlor)
CASRN	72-54-8	50-29-3	72-55-9	72-43-5
Structure				·
ChemiDplus similarity score (%)	100	77	67	65
DSSTox similarity score (%)	100	96	61	52

²Analogues represent a set of structurally similar chemicals identified using two publicly available similarity databases (ChemiDplus and DSSTOX) prefiltered on the basis of availability of health reference values for non-cancer oral toxicity from regulatory agencies, including ATSDR (2002a, b) and U.S. EPA (2017 b, c)



Putative Toxicity Targets for p,p'-DDD and Analogues

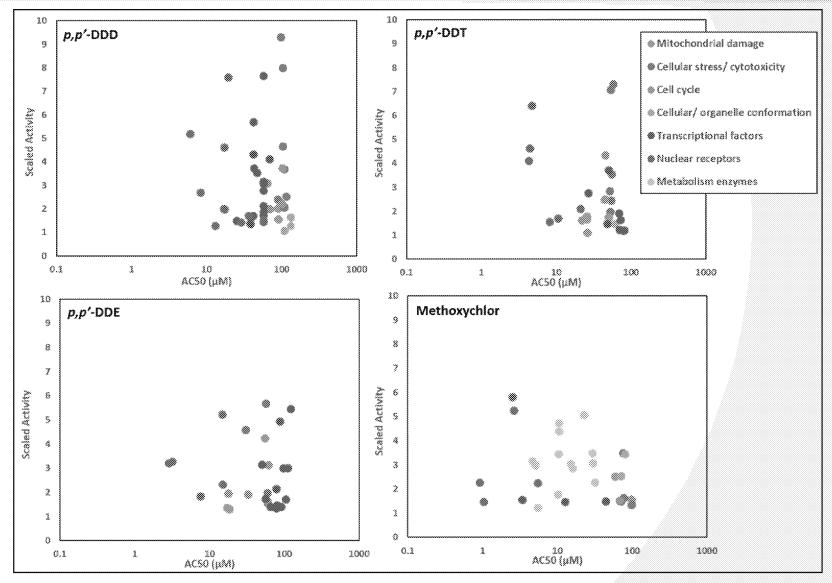
- Liver
- Reproductive





Analysis of Similarities in Bioactivity between p,p'-DDD and Analogues

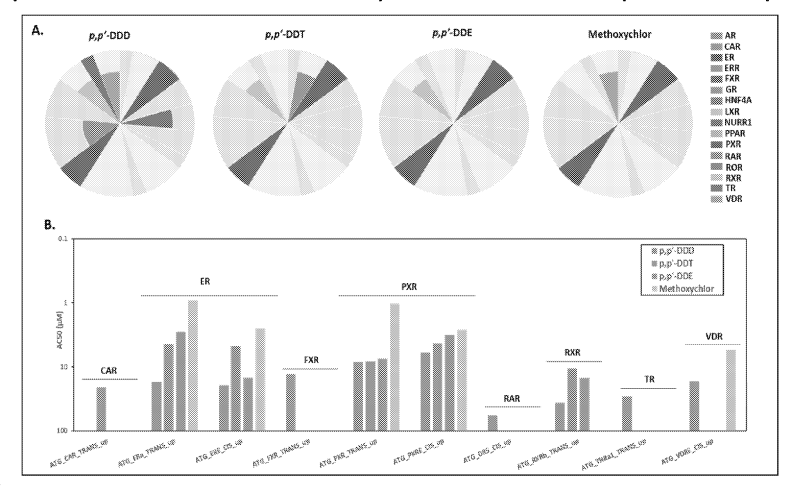
• p,p'-DDD and analogues exhibit similarities in cell-specific responses and target gene pathways in in vitro ToxCast assays conducted in Human Liver Cells





Comparative receptor activation between p,p'-DDD and Analogues

• p,p'-DDD and analogues exhibit similar upregulation of Steroid/Xenobiotic-sensing Nuclear Receptors in *in vitro* ToxCast assays conducted in Hepatoma HepG2 Cells





Summary of Comparative ER/AR Bioactivity

 p,p'-DDD and analogues exhibit similar estrogenic and anti-androgenic activities in *in vitro* ToxCast assays and model predictions for the ER and AR across multiple tissues and cell lines

Table 2, 10	x Cast Clear ivily Summa	ny and Model Predictions	Seores (AND values) (or Ets a	ne AR activities
	p,p'-DDD	p,p'-DDT	p,p'-DDE	Methoxychlor
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elive/Total Assays (A)	7/18 (39)	11/18 (61)	8/18 (44)	14/18 (78)
SSO values (pM)	Range = 14.0 - 32.4	Range = 3.3 - 59.8	Range = 3.5 - 46.2	Range = 0.9 - 44.2
	Median = 18.7	Median = 6.1	Median = 16.5	Median = 4.6
gonisi adilying BC value (95% Ci) ^b	0.0715 (0.0342-0.0738)	0.190 (0.181-0.231)	0.0679 (0.0614-0.0963)	0.254 (0.247-0.260)
ntagonist activity JC value (95% CI)	0	0	0	0
:live/Total Assays (%)	4/11 (36)	AR assays 3/11 (27)	4/11 (36)	3/11 (27)
CSB values (pM)	Range = 31.0 - 62.8	Range = 17.8 - 72.0	Range = 7.0 - 58.7	Range = 29.3 - 40.8
	Median = 44.8	Median = 47.0	Median = 29.6	Median = 34.2
ganist activity IC value (95% Ci)	0	0	0	0
ntagonist activity UC value (95% CI)	0.0973 (0.0649-0.124)	0.0642 (0.0318-0.108)	0.251 (0.234-0.291)	0.0429 (0.0364-0.0465)

⊕ata were sourced from Judson et al. (2015) and Kleinstreuer et al. (2018). ^{p.} 95% CI for the ER activity model were sourced from a subsequent publication to the Judson et al., (2015) study (Watt and Judson, 2018).

Abbreviations: AUC = area under the curve score ranging from 0−1. An AUC value of 0 indicates that the chemical is inactive; CI = confidence interval.

Table 3. Evidence I	Integration Summary and Conclusion		
Similarity Context	Summary of Findings	Evidence Integration conclusions	
Structure and physicochemical properties	 p,p'-DDD and identified analogues (p,p'-DDT and p,p'-DDE and methoxychlor) demonstrate similarities in basic structural features (chlorinated diphenylalkane structure) 	 p,p'-DDT is selected as a suitable source analogue for the assessment of non-cancer oral toxicity of p,p'-DDD based largely on toxicokinetic similarities, with supportive 	
	 p,p'-DDT and p,p'-DDE also share key functional groups (p,p'-chlorine substituents) and physicochemical properties important for bioavailability (lipophilicity and low BCF values) with p,p'-DDD 	information from <i>in vivo</i> toxicity testing, structural similarity evaluations and <i>in vitro</i> bioactivity from HTS assays	
Toxicokinetics	 p,p'-DDT is a metabolic precursor of p,p'-DDD and both chemicals show similarities in toxicokinetics (Absorption, Distribution and Metabolism [ADME]) in humans and experimental animal models (preferential partitioning into fat, similar metabolism and excretion pathways and prolonged elimination rates) 	EF£9998£177800 EN£ 08-20-2017	
	 Other analogues demonstrate differences in ADME in comparison to the target. p,p'-DDE is less metabolically active; methoxychlor is metabolized differently and appears to be less bioaccumulative 	Provisional Peer-Reviewed Toxicity Values for $p_{i}p'$ -Dichlorodiphenyldichloroethane $(p_{i}p'$ -DDD) (CASEN 72-54-8)	
Toxicodynamic	 Consistency and coherence across health effects in experimental animals for non-cancer oral toxicity among the analogues point to putative toxicity targets for p,p'-DDD (primarily liver and reproductive toxicity) 	Superfund Healfn Risk Technical Support Center National Centre for Environmental Assessment Office of Research and Decelopment U.S. Environmental Protection Agency	
	 Similarities in in vitro bioactivity profiles from ToxCast assays between the target and analogues with respect to cell-specific responses and target gene pathways provide mechanistic plausibility for the liver and reproductive effects associated with this group of chemicals 	U.S. Environmental Protection Agency Cracimenti, OH 49368	



Strengths, Limitations, and Challenges of Expert-Driven Read Across

Strengths

- Provides an opportunity to develop human health values where none would be possible based on traditional risk assessment approaches and practice
- Approach is flexible, nimble, and evergreen

Limitations

- Is not high-throughput per se
- Current approach is dependent on analogue space with existent health values

Challenges

- Toxicokinetics (e.g., metabolism) is a critical similarity context that is often highly datapoor
- Increase throughput...



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